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NAVAL POSTGRADUATE SCHOOL

MONTEREY, CALIFORNIA

THESIS

**IMPROVING CLUSTER ANALYSIS WITH AUTOMATIC
VARIABLE SELECTION BASED ON TREES**

by

Anton D. Orr

December 2014

Thesis Advisor:
Second Reader:

Samuel E. Buttrey
Lyn R. Whitaker

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**IMPROVING CLUSTER ANALYSIS WITH AUTOMATIC VARIABLE
SELECTION BASED ON TREES**

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Commander, United States Navy
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Submitted in partial fulfillment of the
requirements for the degree of

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**NAVAL POSTGRADUATE SCHOOL
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ABSTRACT

Clustering is an algorithmic technique that aims to group similar objects together in order to give users better understanding of the underlying structure of their data. It can be thought of as a two-step process. The first step is to measure the distances among the objects to determine how dissimilar they are. The second clustering step takes the dissimilarity measurements and assigns each object to a cluster.

We examine three distance measures proposed by Buttrey at the Joint Statistical Meeting in Seattle, August 2006 based on classification and regression trees to address problems with determining dissimilarity. Current algorithms do not simultaneously address the issues of automatic variable selection, independence from variable scaling, resistance to monotonic transformation and datasets of mixed variable types.

These “tree distances” are compared with an existing dissimilarity algorithm and two newer methods using four well-known datasets. These datasets contain numeric, categorical and mixed variable types. In addition, noise variables are added to test the ability of each algorithm to automatically select important variables. The tree distances offer much improvement for the problems they aimed to address, performing well against competitors amongst numerical datasets, and outperforming in the cases of categorical and mixed variable type datasets.

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TABLE OF CONTENTS

I.	INTRODUCTION.....	1
II.	BACKGROUND AND LITERATURE REVIEW	5
A.	DISSIMILARITY MEASUREMENTS FOR NUMERIC VARIABLES	6
1.	Euclidean and Manhattan Distance.....	6
2.	Variable Weights.....	8
B.	DISSIMILARITY MEASUREMENTS FOR MIXED VARIABLES	10
1.	Binary Variables.....	10
2.	General Categorical Variables.....	12
3.	Mixed Variables.....	13
C.	CLASSIFICATION AND REGRESSION TREES.....	14
1.	Regression Trees.....	14
2.	Classification Trees.....	15
3.	Random Forest Distance.....	16
D.	CLUSTERING ALGORITHMS	18
1.	Partitioning Algorithms.....	18
a.	<i>K-means</i>	18
b.	<i>PAM</i>	19
2.	Hierarchical Algorithms	19
3.	Sparse Clustering.....	20
a.	<i>Sparse K-means</i>	20
b.	<i>Sparse Hierarchical</i>	20
III.	METHODOLOGY.....	23
A.	INTRODUCTION.....	23
B.	BUTTREY TREE DISTANCES	23
1.	Tree Distance 1, d_1	24
2.	Tree Distance 2, d_2	24
3.	Tree Distance 3, d_3	25
C.	ALGORITHMS USED	26
D.	EVALUATION USING CRAMÉR'S V	27
E.	DATASETS	27
1.	Iris	28
2.	Optical	29
3.	Splice	29
4.	Cardiac Arrhythmia	29
IV.	ANALYSIS.....	31
A.	INTRODUCTION.....	31
B.	RESULTS.....	31
C.	CONCLUSION	34
V.	SUMMARY.....	35
A.	SUMMARY	35

B. FUTURE WORK.....	35
LIST OF REFERENCES.....	37
INITIAL DISTRIBUTION LIST	39

LIST OF FIGURES

Figure 1.	Illustration of Euclidean distance.....	7
Figure 2.	Illustration of Manhattan distance.....	8
Figure 3.	Tree with deviance in large circles and leaf numbers in small circles (from Buttrey, 2006)	25
Figure 4.	Example tree evaluated using d_3 with deviances for leaf 14 and 15 (from Lynch, 2014)	26

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LIST OF TABLES

Table 1.	Observation i and j association table.....	10
Table 2.	Dimensions of datasets	28
Table 3.	Iris Dataset Results	32
Table 4.	Optical Digits Dataset Results.....	33
Table 5.	Splice Dataset Results	33
Table 6.	Cardiac Arrhythmia Dataset Results	34

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LIST OF ACRONYMS AND ABBREVIATIONS

Agnes	agglomerative nesting
BCSS	between cluster sums of squares
CART	classification and regression trees
Daisy	DISsimilAritY
PAM	partitioning around medoids
PMA	penalized multivariate analysis
SPC	sparse principal components
UPGMA	unweighted pair-group average method
WCSS	within cluster sums of squares

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EXECUTIVE SUMMARY

Grouping similar objects together is an instinctual response when trying to organize or process an array of information. This sorting and classification process dates back to Aristotle's work on biological taxonomy (Hartigan, 1975). Since then, most fields have developed their own processes to accomplish the task, and data scientists use clustering algorithms.

Clustering can be thought of as a two-step process. The first step is the measurement of distance, or more generally, dissimilarity between observations. The second step is assigning each object to a specific group or "cluster." Our focus is to improve clustering techniques that are applied to the types of data encountered by military practitioners. Often such data consists of a mix of both numeric and categorical data. For example, military manpower studies of promotion rates include numeric variables such as "age" and "number of dependents" as well as categorical variables such as "education level," "commissioning source" and "marital status" (Kizilkaya, 2004). In addition, it is common that in such studies the number of variables is great, but the number of relevant variables for clustering is small.

The book by Kaufman and Rousseeuw (1990) presents different variable types and ways to measure distance along with the two main clustering methods. Numerical variables can be measured using either Euclidean or Manhattan distance, although both are susceptible to scaling or monotonic transformation. Dissimilarities for categorical and mixed data types can be measured using a generalization of Gower's (1971) formula. A common practice is to use Euclidean distance when all variables are numeric and Gower dissimilarity for datasets of mixed variable types as is implemented by the daisy function in the R software (R Core Team, 2014) package "cluster" (Maechler, Rousseeuw, Struyf, Hubert, & Hornik, 2014).

Current clustering algorithms do not simultaneously address these key issues:

- Independence from variable scaling
- Automatic selection of important variables
- Resistance to monotonic transformation
- Inputs with mixed numerical and categorical variables.

A new approach by Buttrey (2006) uses classification and regression trees to define three new distances (tree distances) that attempt to address these issues. This thesis compares these tree distances to dissimilarities computed by daisy and the random forest proximities proposed by Breiman (2001). These five distances are computed for four existing and well-known datasets and input into the clustering algorithms Agnes (agglomerative nesting), PAM (partitioning around medoids) and K -means. Noise is introduced into each dataset to test automatic variable selection. An additional method that uses sparse hierarchical and sparse k -means combines the distance measurement and clustering steps is introduced by Witten and Tibshirani (2010) and is also used for comparison.

The tree distances take a dataset of n observations and p variables and create a classification or regression tree using each variable as the response. After the trees are created, each object is run down the tree. For the first distance d_1 , two objects are considered similar if they end up in the same leaf. The second distance, d_2 , scales the dissimilarity of objects by the “quality” of each tree, which is determined by the amount of deviance between the root node and leaves. The third distance, d_3 , computes dissimilarity between objects by comparing deviance at their parent node to the deviance at the root node.

Of the distance metrics tested, d_2 consistently performed well across all four datasets.

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I. INTRODUCTION

Classification dates back to Aristotle's biological work on the taxonomy of plants and animals. Clustering is the grouping of similar objects and is a counterpart of classification (Hartigan, 1975). Algorithms, whose modern usage has extended into the fields of astronomy, medicine, economics, and many other types of data analysis, implement the more formal technique of clustering. Military and industry practitioners of data analytics face the challenge of turning an abundance of information into useful tools that can assist in making critical decisions. The amount of data collected by private industry and government institutions can be on the order of terabytes daily and clustering techniques can aid the decision-making process by helping to identify and visualize the patterns in this data. Finding the signal in this sea of "noise" will require improvements to the tools we currently use.

Our focus is to improve clustering techniques that are applied to the types of data encountered by military practitioners. Often such data consists of a mix of both numeric and categorical data. For example, military manpower studies of promotion rates include numeric variables such as "age" and "number of dependents" as well as categorical variables such as "education level," "commissioning source," and "marital status" (Kizilkaya, 2004). In addition, it is common that in such studies the number of variables is great, but the number of relevant variables for clustering is small. In this thesis, we build on the work of Lynch's (2014) empirical study of a new and novel clustering technique from Buttrey (2006), specifically designed to cluster observations with mixed categorical and numeric variables and which promises to be robust when applied to datasets with a large number of irrelevant and noisy variables.

Clustering is an algorithmic technique. Which algorithm is used depends on a number of factors including the tradition of the field of application, the types of variables measured, the size of the data set, and what is known a priori about the clusters. Most research in this area has been for data consisting solely of

numeric variables (e.g. Ooi, 2002), although there are notable attempts to treat clustering of categorical variables and mixed variables (e.g. Zhang, Wang and Zhang, 2007). A recent research trend in the area of clustering concerns variable selection, i.e., when confronted with a large (big) data set with many variables, which should be used or weighted more heavily when clustering (e.g. Witten & Tibshirani, 2010). Few works explicitly address both the issues of clustering mixed variables and variable selection.

The process of clustering can be thought of as a two-step process. At the center is the measurement of distance, or more generally, dissimilarity between observations. This aspect of clustering is given much less attention in the literature than the second step (Hastie, Tibshirani and Friedman, 2009). The second step, the clustering step, is the process of grouping like objects into clusters. For clustering based on numeric variables, the choice of dissimilarity measurement is problematic. It depends, among other things, on the choice of an appropriate scale for each variable and on how much weight each variable will be given in the dissimilarity computation. Combining categorical variables with numeric variables complicates the choice of dissimilarity measurement further.

Current clustering algorithms that implicitly or explicitly produce distance calculations do not simultaneously address these key issues:

- Independence from variable scaling
- Automatic selection of important variables
- Resistance to monotonic transformation
- Inputs with mixed numerical and categorical variables

The fundamental contribution of Buttrey (2006) is his approach to using classification and regression trees to define distances (tree distances) that address each of these four key issues. Lynch (2014) studies three tree distances and shows that clustering based on two of the tree distances is advantageous when clustering noisy data of mixed variable types. But Lynch (2014) does not completely decouple the process of measuring dissimilarity from the process of

clustering. In our work, we explicitly decouple the dissimilarity measuring process from the clustering process. This allows us to include in our study a comparison of tree distances with two competing dissimilarities for mixed variables. By decoupling the dissimilarity computations from the clustering algorithm, we also extend the sparse clustering methods of Witten and Tibshirani (2010) so that they can be used to cluster mixed variable datasets. This allows us to experiment with the use of more modern clustering algorithms that are not explicitly designed for mixed variables.

This thesis examines the ability of five methods for determining dissimilarities amongst observations in datasets with and without mixed variables. Their performance will be evaluated based on the clustering solutions they produce for four existing and well-known datasets. The five dissimilarities include the three new tree distances and two existing dissimilarities: one of Euclidean distance for numeric variables or the ubiquitous Gower (1971) dissimilarity, a generalization of the Gower coefficient, for mixed variables, and the lesser known distance based on Breiman's (2001) random forest proximities. Each will be matched with at least two clustering algorithms to establish whether the new tree distances produce better results, and if so, what the causes are. In addition, because a benefit of tree distances is their robustness when noisy or irrelevant variables are present, we also produce results for two clustering methods of Witten and Tibshirani (2010). Their sparse clustering incorporates variable selection into the distance computations and has been observed to perform well for datasets with few relevant variables.

This paper is organized as follows. Chapter II introduces common dissimilarities for numeric, categorical, and mixed variables and describes distances based on random forest proximities. Chapter II also reviews some of the current clustering methodology. Chapter III is an introduction to the new tree distances and review of the datasets used in the study. Chapter IV presents findings and analysis of the research. Finally, chapter V provides conclusions and discusses future work to be done with these new distances.

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II. BACKGROUND AND LITERATURE REVIEW

There are many approaches to the problem of clustering and many finely detailed aspects addressed in academic texts and papers. Two often referenced sources on which we rely heavily to lay the groundwork for clustering are the books by Kaufman and Rousseeuw (1990) and Hastie, Tibshirani and Friedman (2009). Kaufman and Rousseeuw (1990) cover many of the algorithms used to measure dissimilarity between observations and group them into clusters. The second book by Hastie, Tibshirani and Friedman (2009), covers many of the same techniques as do Kaufman and Rousseeuw (1990) and introduces a unique distance measurement technique that will be examined further in this paper.

The basic idea for clustering is to take a dataset with n observations and p measurements (variables) and separate them into groups, or “clusters,” based on their proximities. Thus, in this chapter, we begin by discussing common methods for measuring proximities starting with numeric variables and then discussing mixed numeric and categorical variables. We then give a review of classification and regression trees (CART). CART forms the basis of the new tree distances introduced in Chapter III. CART is also the basis of a competing distance computed from the Breiman (2001) random forest proximities and described in this chapter.

We note that most clustering algorithms do not require that a distance characterize proximities between observations. Most clustering algorithms will take dissimilarities rather than distances, where the dissimilarities have the following properties: the dissimilarity of an object with itself is zero, all dissimilarities are non-negative, and the dissimilarity between object i and object j is the same as the dissimilarity between object j and object i .

The difference between a dissimilarity and a distance is that the triangle inequality need not hold for dissimilarity. Throughout the thesis, we use the less

restrictive term dissimilarity to mean either distance or dissimilarity. Further, proximity is the general term which can be measured by how far apart objects are or by closeness or similarity, where similarity is a monotonic decreasing function of dissimilarity (often similarity is one minus dissimilarity).

A. DISSIMILARITY MEASUREMENTS FOR NUMERIC VARIABLES

There are many ways to measure the dissimilarity between objects i and j . Two of the most common for continuous numeric variables are Euclidian distance and Manhattan distance. These are suitable for variables that are positive or negative real numbers on a linear scale, such as age, weight, cost, length, etc. Numeric variables can be subjected to a change in units or monotonic transformations that will affect their distance. For linear forms, the primary way to handle these changes is standardization, so that relative distances between observations remain constant.

1. Euclidean and Manhattan Distance

Euclidean distance is the true geometrical distance between two points i and j with coordinates (x_{i1}, \dots, x_{ip}) and (x_{j1}, \dots, x_{jp}) , and is given by the equation:

$$d(i, j) = \sqrt{(x_{i1} - x_{j1})^2 + (x_{i2} - x_{j2})^2 + \dots + (x_{ip} - x_{jp})^2} \quad (2.1)$$

where the x 's are numeric and the number of variables are measured by p (Kaufman and Rousseeuw, 1990). Equation 2.1 gives the length of the hypotenuse of the triangle, so this distance is sometimes referred to as Pythagorean distance and is illustrated in Figure 1.

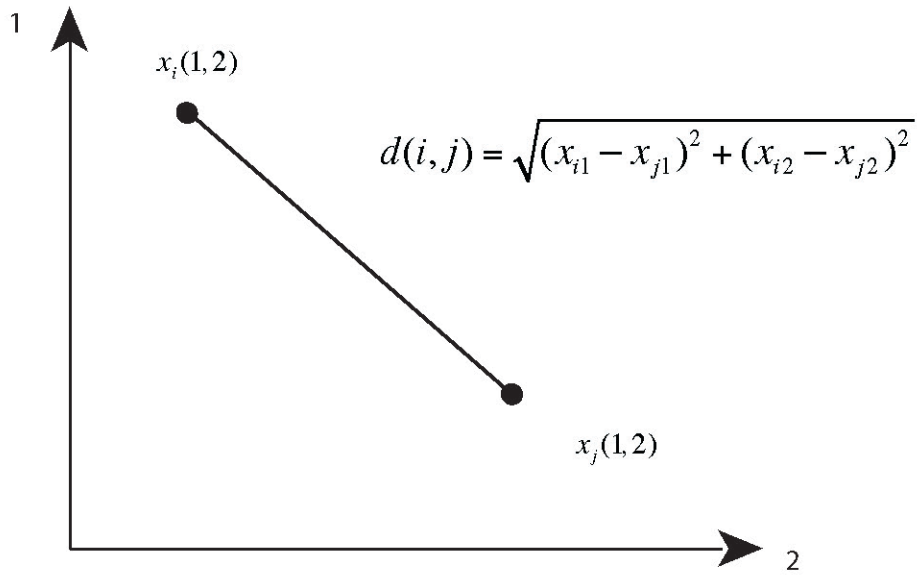


Figure 1. Illustration of Euclidean distance.

The second well-known distance metric is the *Manhattan distance*, or *city block*. This is the distance between two points in a grid based strictly on a horizontal and/or vertical line of travel instead of the diagonal route. This distance is the sum of the absolute value of the difference between each variable:

$$d(i,j) = |x_{i1} - x_{j1}| + |x_{i2} - x_{j2}| + \dots + |x_{ip} - x_{jp}| \quad (2.2)$$

and is illustrated in Figure 2.

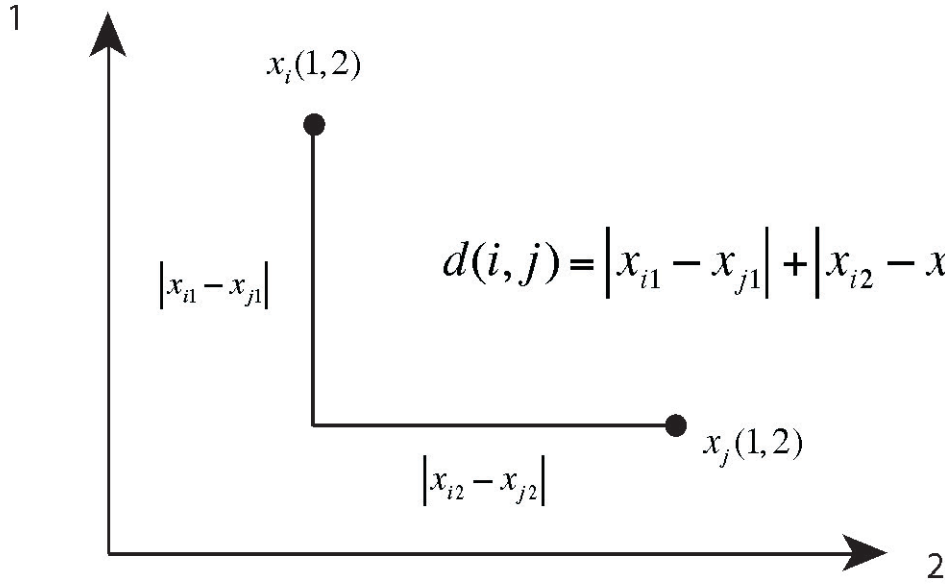


Figure 2. Illustration of Manhattan distance.

The Minkowski distance provides a generalization of these distances

$$d(i, j) = \left(|x_{i1} - x_{j1}|^q + |x_{i2} - x_{j2}|^q + \dots + |x_{ip} - x_{jp}|^q \right)^{1/q} \quad (2.3)$$

in which q is any real number greater than or equal to 1 (Kaufman & Rousseeuw, 1990).

The distance algorithm daisy (DISssimilarity) implemented in the R package (R Core Team, 2014) “cluster” (Maechler, Rousseeuw, Struyf, Hubert, & Hornik, 2014), which will be discussed later and which we use in this study, can use either the Euclidean or Manhattan distance for its calculations when all variables are numeric.

2. Variable Weights

The distances (2.1), (2.2), and (2.3) weigh each variable equally. Other considerations in distance measurements concern the importance of individual variables and their weight towards the calculations. For example, changing the scale of a variable by converting from meters to kilometers would change the apparent weight of that variable by a factor of 1000. One way to deal with this is

through standardization of the variables. Kaufman and Rousseeuw (1990) accomplish this with three steps. First, calculate the mean for variable f ,

$$m_f = \frac{1}{n} (x_{1f} + x_{2f} + \dots + x_{nf}) . \quad (2.4)$$

Then take the *mean absolute deviation*

$$s_f = \frac{1}{n} \{ |x_{1f} - m_f| + |x_{2f} - m_f| + \dots + |x_{nf} - m_f| \} . \quad (2.5)$$

Use the mean (2.4) and the mean absolute deviation (2.5) to standardize the variable in a “z-score” calculation

$$z_{if} = \frac{x_{if} - m_f}{s_f} . \quad (2.6)$$

Standardization (2.6) gives all variables an equal weight independent of their units (Kaufman & Rousseeuw, 1990). Often practitioners use either the sample standard deviation or the range in place of the mean absolute deviation in (2.5). The advantage of using the mean absolute deviation over the sample standard deviation or the variable range is that it is more robust to extreme values.

More generally, it must be decided if all variables should have equal importance before clustering. Some variables matter more than others and this consideration requires some subject matter expertise. A hypothetical dataset of aircraft might include age, top speed, gross weight, empty weight, cost and tail number. If you wanted to cluster the aircraft according to cargo-carrying capability, gross weight might be a better discriminant than age, and should be weighted appropriately. Tail number probably provides no useful information, so it might be excluded or equivalently given a weight of 0. Weighted Euclidian distance is given by

$$d(i, j) = \sqrt{w_1(x_{i1} - x_{j1})^2 + w_2(x_{i2} - x_{j2})^2 + \dots + w_p(x_{ip} - x_{jp})^2} \quad (2.7)$$

where w is the weight applied to each variable, and can be used to indicate perceived importance (Kaufman & Rousseeuw, 1990). Both Kaufman and Rousseeuw (1990) and Hastie, Tibshirani and Friedman (2009) see this process as subjective and therefore cumbersome in data with many variables. However,

current work (Witten and Tibshirani, 2010) seeks to automate this process through variable selection, which computes the variable weight, with selected (irrelevant) variables getting weight zero.

B. DISSIMILARITY MEASUREMENTS FOR MIXED VARIABLES

Different types of variables require different approaches when considering their proximities for clustering. For non-numeric variables, we start by considering dissimilarities based only on categorical variables with two levels each (binary variables). We then discuss dissimilarities between nominal and ordinal categorical variables with two or more levels and then show the most common approach to combining numeric and categorical variables.

1. Binary Variables

Categorical variables with two levels are common in practice. For example, an attribute can be present/absent, yes/no, male/female, or small/large. Often such variables are encoded as binary 0/1 variables. When these characterizations are symmetric, of equal value, similar weight, and all variables in a dataset are binary, the similarity between two observations is calculated with a matching coefficient derived from an association table:

Table 1. Observation i and j association table

		object j		
		1	0	
object i	1	a	b	$a+b$
	0	c	d	$c+d$
		$a+c$	$b+d$	

In Table 1, the value depicted by a is the sum of binary attributes where object i and object j both equal 1 and d is where they both equal 0. The matching coefficient for similarity $s(i, j)$ is then computed as

$$s(i, j) = \frac{a + d}{a + b + c + d} , \quad (2.8)$$

where the numerator is the total number of matches and the denominator, the total number of binary variables (Kaufman & Rousseeuw, 1990).

The corresponding dissimilarity would be $d(i, j) = 1 - s(i, j)$, with coefficients taking on values from 0 to 1. The algorithm daisy default is to use this coefficient to compute dissimilarities between observations when all variables are binary categorical variables.

Another type of binary variable is asymmetric. One value, usually taken to be 1, of an asymmetric binary variable is rare, so most objects would have a value of 0. When compared for similarity, negative matches would be expected, so positive matches are considered more significant.

An example dataset of aircraft might include a variable of aircraft type, measuring whether the aircraft is an attack aircraft, with value 1, or not, with value 0. In this case, for variable f corresponding to aircraft type the statement $x_{if} = 1$ and $x_{jf} = 1$ is a rare positive match, and implies that aircraft i and j are both attack aircraft. In this dataset of civilian aircraft, a negative match “non-attack,” is common and less meaningful because the two could be of many different aircraft types. To calculate similarities, only positive matches are considered, where $s_{ijf} = 1$ for each match at variable f , giving similarity between aircraft i and j

$$\frac{\sum_{f=1}^p s_{ijf}}{p} . \quad (2.9)$$

To accommodate missing values, let $s_{ijf} = 1$ if variable f can be measured for both object i and j and define the similarity to be

$$s(i, j) = \frac{\sum_{f=1}^p \delta_{ijf} s_{ijf}}{\sum_{f=1}^p \delta_{ijf}} . \quad (2.10)$$

This is Gower's (1971) general similarity coefficient for asymmetric binary categorical variables, and can be implemented in daisy.

2. General Categorical Variables

The last variable type is categorical. This variable comes in the form of nominal or ordinal values. Nominal variable types are for observations that take on multiple values, where the values are not ranked and are simply labels. Ordinal values have a ranking attached.

An example of a nominal categorical variable would be the county in which a person resides, with the counties encoded by values one through five. When measuring dissimilarity, the difference between counties one and two should be the same as to the difference between counties two and five. The most common way to calculate dissimilarity for a nominal categorical variable f is a simple matching approach:

$$d_{ijf} = \begin{cases} 0 & \text{if } x_{if} = x_{jf} \\ 1 & \text{if } x_{if} \neq x_{jf} \end{cases} . \quad (2.11)$$

When all p variables are nominal categorical variables the dissimilarity is calculated as the average of individual dissimilarities that are available for both objects i and j :

$$d(i, j) = \frac{\sum_{f=1}^p \delta_{ijf} d_{ijf}}{\sum_{f=1}^p \delta_{ijf}} . \quad (2.12)$$

Ordinal variables take on different states like nominal values, except they have a meaningful order. An example would be responses to survey questions with values one through five on a five point Likert scale. The states range from strongly disagree = 1, to strongly agree = 5, and the difference between states 1 and 2 would not be equivalent to the difference between 2 and 5.

Dissimilarities between objects that contain ordinal variables usually assign a score to each value of the ordinal variable. A data-driven approach to

scoring is to base the scores on the ranks of the observations. For example, if the ordered states are 1, 2,..., M , a *z-score* can be created using the rank r_{if} of the i -th observation in the f -th variable by

$$z_{if} = \frac{r_{if} - 1}{M_f - 1} \quad (2.13)$$

where M_f is the highest state for variable f . The *z-score* will take on values 0 to 1. When all p variables are ordinal, the dissimilarity between observations i and j can then be calculated by the Manhattan distance divided by the number of variables being considered (Kaufman & Rousseeuw, 1990).

3. Mixed Variables

So far, we are able to compute distances for five variable types: numeric, symmetric and asymmetric binary, and nominal and ordinal categorical. When different types occur in the same dataset, we need a way to combine them. In this section we describe Gower's approach to computing dissimilarities for mixed variables. This approach is implemented by daisy as the default for datasets with mixed variables.

First, the dissimilarity for each variable is scaled to be between 0 and 1, i.e., for any pair of objects i, j the dissimilarity for variable $f = 1, \dots, p$ is $0 \leq d_{iff} \leq 1$. Once these calculations have been made, an overall dissimilarity can be defined for datasets that contain p variables of a mixed nature

$$d(i, j) = \frac{\sum_{f=1}^p \delta_{iff} d_{iff}}{\sum_{f=1}^p \delta_{iff}}, \quad (2.14)$$

where, as before, δ_{iff} is equal to 1 when both x_{if} and x_{jf} for the f -th variable are non-missing (Kaufman & Rousseeuw, 1990).

In (2.14) d_{iff} is the contribution of the f -th variable to the dissimilarity between i and j . If either observation is missing on variable f , $\delta_{iff} = 0$ and that variable does not contribute to the dissimilarity. For binary or nominal variables,

(2.10) gives the dissimilarity. For continuous or ordinal variables, the dissimilarity is given by

$$d_{iff} = \frac{|x_{if} - x_{jf}|}{R_f} \quad (2.15)$$

where R_f is the range of variable f ,

$$R_f = \max_h x_{hf} - \min_h x_{hf} \quad (2.16)$$

and h covers all non-missing observations for variable f .

C. CLASSIFICATION AND REGRESSION TREES

When dealing with datasets that involve a large number of variables, classification and regression trees (CART) are often used (Breiman, Friedman, Stone, & Olshen, 1984). These are simple nonlinear predictive models that recursively partition a data space and fit a decision tree into each partition. Regression trees are used when the response variable has continuous values, and classification trees when a response has a finite number of unordered values or “classes.”

1. Regression Trees

Regression tree algorithms take an observation with a numeric response y and p input variables for each of n observations: that is $(y_i | x_i)$ for $i = 1, 2, \dots, n$, with input variables $x_i = (x_{i1}, x_{i2}, \dots, x_{ip})$ (Hastie, Tibshirani, & Friedman, 2009). The algorithm automatically decides the splitting variable and split to produce a partition with M regions R_1, R_2, \dots, R_M . For each region, the predicted response, $y(x)$ is modeled as a constant c_m , so that

$$y(x) = \sum_{m=1}^M c_m I(x \in R_m) \quad (2.17)$$

where $I(x \in R_m)$ is the indicator function which is 1 if $x \in R_m$ and 0 otherwise. Using minimized sum of squared deviations between predicted and observed response as a criterion for selecting c_m , the best \hat{c}_m for a numeric y is the average of y_i in region R_m

$$\hat{c}_m = \text{ave}(y_i | x_i \in R_m) . \quad (2.18)$$

A tree is grown starting with the root node, which contains all observations. Proceeding with a greedy algorithm, at each node R , a splitting variable j and split point s (or set s for categorical *variables*) will partition the node R into left and right child nodes defined by

$$R_L(j,s) = \{x_{ij} \in R | x_{ij} \leq s\} \text{ and } R_R(j,s) = \{x_{ij} \in R | x_{ij} > s\} \quad (2.19)$$

for numeric variables x_j . For categorical variables,

$$R_L(j,s) = \{x_{ij} \in R | x_{ij} \in s\} \text{ and } R_R(j,s) = \{x_{ij} \in R | x_{ij} \notin s\} \quad (2.20)$$

where s is a subset of the levels of the categorical variable.

The values for j , s and c_L , c_R , the predicted values for y in the left and right child nodes respectively are found by solving

$$\min_{j,s} \left[\min_{c_L} \sum_{x_{ij} \in R_L(j,s)} (y_i - c_L)^2 + \min_{c_R} \sum_{x_{ij} \in R_R(j,s)} (y_i - c_R)^2 \right] , \quad (2.21)$$

the combined “impurity” of the two child nodes (Hastie, Tibshirani, & Friedman, 2009).

For each splitting variable j and split s , the inner minimization is solved by

$$\hat{c}_L = \text{ave}(y_i | x_i \in R_L(j,s)) \text{ and } \hat{c}_R = \text{ave}(y_i | x_i \in R_R(j,s)) \quad (2.22)$$

This algorithm will grow a tree and stops when a node size limit or a threshold value for equation (2.21) is reached. This tends to produce a tree which is too large. With large trees there is a danger of over-fitting. That is, the tree is grown to a depth that predicts each observation in the data set well, but it cannot predict observations taken from an independent, but similar data set. To guard against over-fitting, the tree is then “pruned” by collapsing internal nodes to find a subtree that minimizes ten-fold cross-validated measure of impurity.

2. Classification Trees

Classification trees are used when the objective is to find which class an object belongs to. A process similar to the regression algorithm is used, with the

main difference being the criteria used to split nodes. The proportion of objects in node m , which represents region R_m and n_m observations is shown by,

$$\hat{p}_{mk} = \frac{1}{n_m} \sum_{x_i \in R_m} I(y_i = k) \quad (2.23)$$

where k is the class. The majority of observations residing in node m determine its classification: $k(m) = \arg \max_k \hat{p}_{mk}$. At each split, a variable and splitting criteria are chosen to minimize the impurity of the child nodes. A common measure of impurity and the one we use is the multinomial deviance,

$$-2 \sum_{k=1}^K n_{mk} \log(\hat{p}_{mk}) \quad (2.24)$$

As with regression trees, classification trees are pruned so that the ten-fold cross-validated deviance is minimized.

3. Random Forest Distance

The random forest concept was introduced by Breiman (2001) and builds multiple classification or regression trees to reduce variance, creating a “forest” of trees. In addition to prediction, Breiman (2003) discusses a number of applications of random forests including the notion of using random forests to measure the proximity between two observations. In this thesis, random forests are used to measure dissimilarities between two observations based on Breiman’s proximities.

The steps for building a random forest based on a dataset with n observations are:

For $b = 1$ to B :

1. Draw n cases at random, with replacement, from the dataset to create a training set. This training set is called a “bootstrapped” sample.
2. Grow tree T_b using the bootstrapped sample by repeating the following steps at each node, until a minimum node size n_{\min} is reached:

a) Select $m \ll p$ variables at random from the original variables.
(typically $m = \sqrt{p}$)

b) Pick the best variable and split among m .

c) Split the node into two child nodes.

The output is the set of trees $\{T_b\}_1^B$.

For a categorical response, the class prediction of the b -th random forest tree will be $\hat{C}_b(x)$, making the final prediction $\hat{C}_{rf}^B(x) = \text{majority vote } \{\hat{C}_b(x)\}_1^B$ (Hastie, Tibshirani, & Friedman, 2009). For a numeric response, $y_b(x)$ is the prediction for the b -th random forest tree and the final prediction is the forest average.

Breiman (2001) uses random forests to measure proximity between observations. In this setting there is no response variable. His solution is to label all n observations in the dataset to have value $y=1$ and then to simulate “observations” for the $y=0$ class. The $y=0$ observations are generated so that their marginal distributions match the marginal distributions of the original data, but so that the variables of the simulated observations are independent. Building random forests to correctly classify the combined data can uncover structural properties, like clusters, of the original $y=1$ observations, which are not present in the simulated $y=0$ observations. Breiman calls this method unsupervised random forests.

After each tree in the random forest is built, all the observations are run down each tree and if two (i and j) end up in the same leaf, their similarity $s(i, j)$ is increased by one. After all similarities are totaled, they are normalized by dividing by the number of trees. This is used to create a distance between any two observations $d(i, j) = 1 - s(i, j)$; i.e., the distance between observations i and j is the proportion of trees for which i and j fall in different leaves.

D. CLUSTERING ALGORITHMS

There are two primary families of clustering methods, implemented by either partitioning or hierarchical algorithms (a third approach, model based clustering, is not discussed here). They both require a dissimilarity matrix as the input. A dissimilarity matrix for n observations is the $n \times n$ matrix of dissimilarities or distances $\{d(i, j)\}^{n \times n}$. The two types of clustering methods (partitioning and hierarchical methods) are often used for comparison to see which produces a better cluster organization.

1. Partitioning Algorithms

Partitioning algorithms take all n observations and group them into K clusters. The user inputs the value for K , so it is advisable to try different values for K to see which output produces the best result. There are many algorithms, but this thesis will discuss the two most popular, K -means and PAM.

a. K -means

Given a set of multivariate observations (x_1, x_2, \dots, x_n) , K -means partitions the n observations into K clusters $C = \{C_1, C_2, \dots, C_K\}$ so as to minimize the within-cluster sum of squares (WCSS). The objective is to find:

$$\arg \min_C \sum_{k=1}^K \sum_{x \in C_k} \|x - \mu_k\|^2 \quad (2.25)$$

where μ_k is the mean of observations in C_k (MacQueen, 1967).

The algorithm proceeds in two steps. First, an initial set of K means m_1, \dots, m_K is chosen at random and each observation is assigned to the cluster that is closest. After all observations have been assigned, the centroid of each cluster becomes the new mean. The process starts again by reassigning observations to the new means until convergence and the assignments stop changing. This does not guarantee a global optimum, so the algorithm can be run multiple times with different randomly selected initial clusters to determine the final result.

K -means uses Euclidean distance in (2.25) to measure the distance from observations in a cluster to the cluster center. In addition, a cluster center is the average of the observations in that cluster. For these two reasons, K -means can only be used when all variables are numeric and Euclidean distance is appropriate.

b. PAM

Partitioning around medoids uses the K -medoids algorithm which is related to K -means. They both attempt to minimize squared error, with the difference being that PAM always chooses a datapoint as the center of a cluster instead of using the centroid of i -th cluster. This algorithm can be more robust to noise and outliers than K -means. Further, PAM can be adapted to handle datasets with mixed variable types. Using the medoid as the cluster center requires no averaging of observations and because a cluster medoid is one of the observations in the dataset, dissimilarity between it and the members of the cluster can be measured by any dissimilarity or distance defined for mixed variables.

2. Hierarchical Algorithms

Hierarchical algorithms do not construct a single partition with K clusters, but offer all combinations of clusters from $K = 1$ (all observations in one cluster) to $K = n$ (each observation in its own cluster). This paper will address the agglomerative nesting (Agnes) technique that starts with $K = n$.

This method begins by joining the two observations that are closest together, leaving $n - 1$ clusters. Once the first cluster is created, for subsequent pairings, the distances are measured using the unweighted pair-group average method (UPGMA). This method measures dissimilarities between all objects in two clusters and takes the average value. This process is continued until $K = 1$. Different techniques are then applied to determine the optimal number of clusters from this output. We note that Agnes can cluster datasets of mixed variable types. Agnes only requires that dissimilarities be computed for all pairs of

observations in the dataset. However, it requires that all $\frac{n(n-1)}{2}$ pairwise dissimilarities be computed, a computational load that can be burdensome for large datasets.

3. **Sparse Clustering**

Clustering data with a large set of p variables introduces the challenge of selecting which ones actually differentiate the observations. Witten, Tibshirani and Hastie (2010) present a method of sparse K -means and sparse hierarchical clustering that address this problem. These two new methods will be used with the two previously described traditional clustering algorithms for this paper's analysis.

a. **Sparse K -means**

The sparse K -means algorithm operates by maximizing the between-cluster (weighted) sum of squares (BCSS), which is the equivalent to K -means minimization of WCSS. A weight w_1, \dots, w_p , is assigned to each of the p variables. The value of $\|\mathbf{w}\| \leq s$, where $\mathbf{w} = (w_1, \dots, w_p)$ and s is a tuning parameter that satisfies $1 \leq s \leq \sqrt{p}$.

The algorithm initializes variable weights as $w_1 = \dots = w_p = \frac{1}{\sqrt{p}}$. It then iterates over the following steps until convergence:

1. Compute C_1, \dots, C_k using standard K -means, while holding \mathbf{w} fixed.
2. Maximize BCSS by adjusting \mathbf{w} .

The final value of \mathbf{w} indicates which variables contributed most to the dissimilarity calculations.

b. **Sparse Hierarchical**

This clustering method uses standard hierarchical clustering algorithms. The difference is a dissimilarity matrix is created by the sparse principal

components (SPC) criterion of Witten, Tibshirani and Hastie (2009) as the input. The methodology for obtaining SPC is implemented in an R package (R Core Team, 2014) called PMA (for penalized multivariate analysis) (Witten, Tibshirani, Gross, & Narasimhan, 2013).

Both sparse K -means and sparse hierarchical clustering algorithms are implemented in an R package called “sparcl” (Witten & Tibshirani, 2013) that will be used in the analysis portion of this thesis. The R implementation of sparse hierarchical clustering in this package is not able to accept categorical variables. We get around this by using daisy to construct a $\frac{n(n-1)}{2} \times p$ numeric matrix of pairwise, component-wise dissimilarities, which the R function will accept. This workaround is only suitable for small datasets, since $\frac{n(n-1)}{2} \times p$ grows quickly with n . The “sparcl” implementation of sparse K -means cannot be made to accept categorical variables.

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III. METHODOLOGY

A. INTRODUCTION

We have decoupled the clustering problem into two phases, determining dissimilarity between observations and using the dissimilarities in an algorithm to find how they are best grouped. The five methods for computing dissimilarity include tree distances, dissimilarities as computed using the daisy algorithm, and random forest proximities. The cluster algorithms are traditional partitioning and hierarchical approaches and their improved sparse methodologies introduced by Witten and Tibshirani (2010).

Our new method addresses the phase one clustering problem of distance calculation using classification and regression trees. This methodology addresses traditional shortcomings with mixed variable datasets, linear and monotonic transformations, and automatic selection of “important” variables.

B. BUTTREY TREE DISTANCES

To build classification and regression trees, all observations start in one node. The first split creates two child nodes and is selected based on which variable minimizes impurities in the new nodes, based on residual sums of squares. This process continues until a minimum node size is reached. This tends to produce a large tree which is then “pruned” with ten-fold cross validation. In this final step, the cross-validation finds the tree size that produces the smallest impurity.

A tree algorithm requires a response variable, and with clustering there are none, so we use each variable as a response and create p trees. There is also a tendency to over-fit, so the grown trees will be “pruned” with ten-fold cross validation. This step will find a tree size that produces the smallest impurity (Buttrey, 2006). Every observation will fall into one leaf of the tree.

1. Tree Distance 1, d_1

The first metric calculates distance between observations based on which leaf they reside in. If two observations fall on the same leaf, their distance is 0 and 1 otherwise. The leaf L an observation i falls into, for tree t , is denoted by $L_t(i)$. Thus

$$d_1(i, j) = \sum_{t=1}^p \left\{ \begin{array}{ll} 0 & \text{if } L_t(i) = L_t(j) \\ 1 & \text{if } L_t(i) \neq L_t(j) \end{array} \right\}. \quad (3.1)$$

These distances create a dissimilarity matrix to be used as the input for a clustering algorithm.

2. Tree Distance 2, d_2

The second metric uses the same calculation for distance between observations. The primary difference is that this approach does not treat all trees equally. Some trees might be “better” than others, and the way this is measured is by the overall decrease in deviance a tree produces. The decrease is based on the difference between the deviance at the root node, D_t , and the sum of deviances of all the leaves. The difference is denoted ΔD_t .

A tree that produces a large ΔD_t is considered to be of better quality. Observations that are on different leaves of this “better” tree are more dissimilar than those on different leaves of a lower-quality tree.

The “best” tree with maximum deviance $\max_t(\Delta D_t)$ will be used to scale our distances in d_2 using this formula:

$$d_2(i, j) = \sum_{t=1}^p \left\{ \begin{array}{ll} 0 & \text{if } L_t(i) = L_t(j) \\ \frac{\Delta D_t}{\max_t(\Delta D_t)} & \text{if } L_t(i) \neq L_t(j) \end{array} \right\}. \quad (3.2)$$

The tree in Figure 3 will be used to illustrate the difference between d_1 and d_2 . The large ovals in this picture are the deviance at that node, and the small

circles are the leaf number. If observations i and j land in leaves 14 and 15, their d_1 distance would be 1.

To calculate the d_2 distance, the numerator value is $\Delta D_t = 10,000 - 4,700$, which is the root node deviance minus the sum of the leaf nodes. From the group of trees built, assume $\max_t(\Delta D_t) = 10,000$. This gives a distance of $(5,300/10,000) = 0.53$. If we were calculating the distance between two leaves on the best tree, it would be $(10,000/10,000) = 1$.

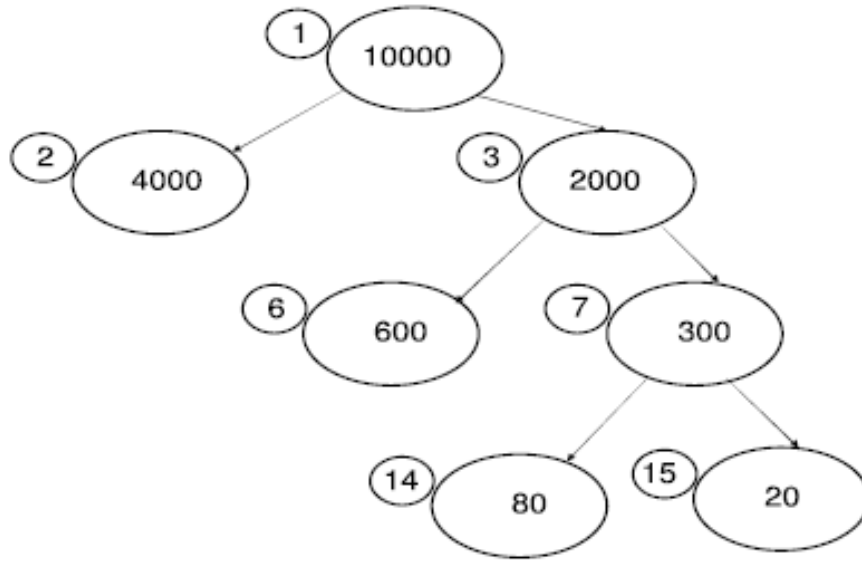


Figure 3. Tree with deviance in large circles and leaf numbers in small circles (from Buttrey, 2006)

3. Tree Distance 3, d_3

The third distance metric compares the deviance at the parent node of two observations to the deviance at the tree's root node. The deviance at the parent is denoted $D_t(i, j)$ and the overall deviance of the tree D_t . The distance is calculated by

$$d_3(i, j) = \begin{cases} 0 & \text{if } L_t(i) = L_t(j) \\ \frac{\Delta D_t(i, j)}{\Delta D_t} & \text{if } L_t(i) \neq L_t(j) \end{cases} \quad (3.3)$$

The process to find the d_3 distance between leaves 14 and 15 is illustrated in Figure 4. Total change in deviance for the tree is $\Delta D_t = 10,000 - 4,700 = 5,300$. After the tree is cropped at the parent node, the change in deviance is $\Delta D_t(14,15) = 4,900 - 4,700 = 200$. The final distance for objects in these leaves is $\frac{\Delta D_t(i,j)}{\Delta D_t} = \frac{200}{5300} = 0.038$. The distance between observations that are close together on a tree are smaller than those that fall farther apart.

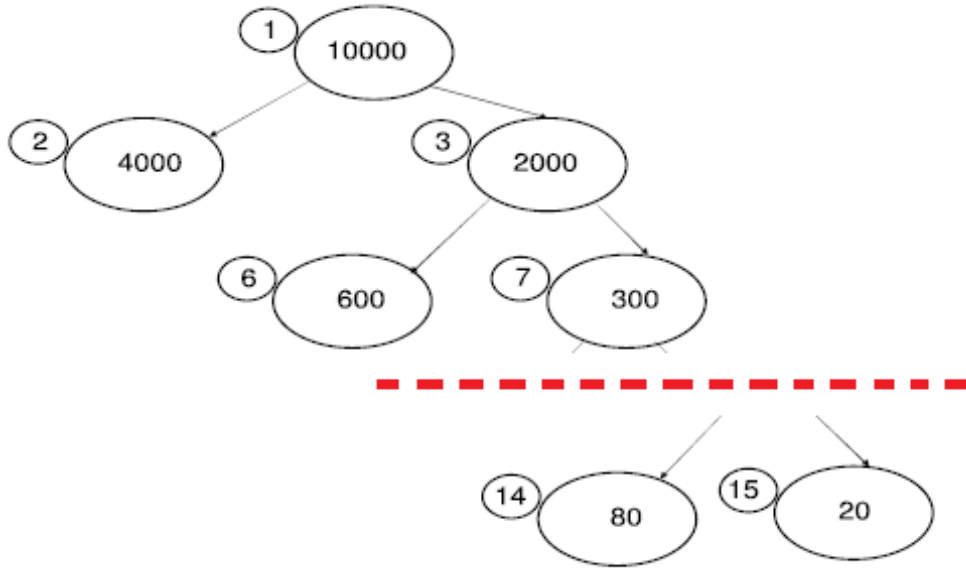


Figure 4. Example tree evaluated using d_3 with deviances for leaf 14 and 15 (from Lynch, 2014)

Currently these three distances are computed and stored for all pairs of observations. However, we need only compute distances among all pairs of leaves within each tree, so substantial gains in efficiency will be possible in the future.

C. ALGORITHMS USED

Five sets of dissimilarity matrices are computed for each dataset. Tree distances are calculated using the steps described. The daisy algorithm

constructs a matrix using Euclidean distance for the numeric datasets and Gower’s dissimilarity for the categorical and mixed datasets. Random forest proximities are used to create the final matrices.

Three clustering algorithms are then used to find a clustering solution, Agnes, PAM and *K*-means. *K*-means clustering is only applied to two of the four datasets, which consist of entirely numeric variables. Because its underlying distance is Euclidean. PAM and Agnes are applied to all four datasets using each of the five dissimilarity measures.

Sparse *K*-means and sparse hierarchical clustering are also used, with sparse *K*-means applied only to the two numerical datasets.

D. EVALUATION USING CRAMÉR’S *V*

To compare different clustering approaches, we used four established datasets in which the “correct” classifications are known. The quality of each solution was evaluated using Cramér’s *V* (Cramér, 1946), which is a scaled version of the χ^2 measure of association for a two-way table. This produces a number between 0 and 1, which will be close to 1 when clusters closely match their class label, and small when they do not.

E. DATASETS

Four datasets from the University of California, Irvine, Machine Learning Repository (Bache & Lichman, 2013) were used in this analysis: Iris, Optical Recognition of Handwritten Digits, Molecular Biology (Splice-junction Gene Sequence), and Cardiac Arrhythmia. The dimensions of each dataset are listed in Table 2. The number of classes *k* is known and will be used to evaluate how well each algorithm classifies a given set of data. In addition, the algorithms will be run with $2k$ clusters because the number of clusters might be greater than the number of classes.

Table 2. Dimensions of datasets

Name	Observations	Variables	Data Type	Classes
Iris	150	4	Numeric	3
Iris + 15 noise	150	19	Numeric	3
Iris + 50 noise	150	54	Numeric	3
Optical	1797	64	Numeric	10
Optical + 15 noise	1797	79	Numeric	10
Optical + 50 noise	1797	114	Numeric	10
Splice	3190	60	Categorical	3
Splice + 15 noise	3190	75	Categorical	3
Splice + 50 noise	3190	110	Categorical	3
Arrhythmia	452	278	Mixed	13
Arrhythmia+ 15 noise	452	293	Mixed	13
Arrhythmia + 50 noise	452	328	Mixed	13

Noise variables are added to see if the algorithm can discern which variables are most important. Therefore, we will introduce 15 and 50 noise variables into each dataset to see how our algorithms perform.

1. Iris

The Iris flower dataset was made famous by Sir Ronald Fisher (Fisher, 1936) and includes information to quantify the variations of iris flowers of three related species. There are 50 samples from each species (Iris setosa, Iris virginica, and Iris versicolor) collected from the Gaspé Peninsula. Measurements were taken of four features of the flowers, the length and width of their sepals and petals in centimeters. Fisher developed a linear discriminant model to distinguish each species from the other and this dataset has been used as a typical test case for many classification techniques. All variables in this dataset are numeric.

Random noise is introduced into the dataset from a normal distribution with a mean of zero and standard deviation of one.

2. Optical

The Optical Recognition of Handwritten Digits dataset was donated by E. Alpaydin and C. Kaynak (Bache & Lichman, 2013). The data is from 43 people who contributed handwritten digits from zero to nine. These were then converted to bitmaps and a preprocessing program was used to determine what number was displayed. The output is an 8 x 8 matrix of 64 integers taking values zero through 16 to account for small distortions in the way different people write their numbers. There are 5,620 observations total, but only 1,797, the original test set, are used in this analysis. All sixty-four variables are numeric.

Noise variables are introduced by a random sample with replacement, of integers zero through 16.

3. Splice

The Splice-junction data comes from Genbank 64.1 (Bache & Lichman, 2013). It includes 3,190 primate DNA sequences with 60 elements. Each element corresponds to a categorical variable with levels C, A, G and T. The purpose of each is to recognize the boundaries between exons (EI) and introns (IE) to determine if the sequence is from a donor or acceptor, respectively. The third class in this dataset is neither. The sixty variables in this dataset are categorical.

The noise variables for splice are generated by drawing from a random sample with replacement, of the letters C, A, G and T, which correspond to DNA sequence elements.

4. Cardiac Arrhythmia

The final dataset comes from H. Altay Guvenir (Bache & Lichman, 2013) of Bilkent University. The variables in the data comprise the electrocardiograph readings from 452 medical patients. The output of these readings contain 279 variables, of which 206 are numeric and 73 categorical. There are 16 response classes with one being normal cardiac function, one undetermined and the rest

denoting differing degrees of cardiac arrhythmia. Only 13 classes are used in the analysis because three of the classes have no patients associated with them.

Noise variables are introduced from a normal distribution with mean of zero and standard deviation of one.

IV. ANALYSIS

A. INTRODUCTION

This chapter presents the results of our comparative analysis of the different dissimilarity measures and clustering algorithms. Cramér's V is the metric used for each algorithm and dataset combination, and a higher value indicates a better pairing, with a value of one being a perfect match between clusters and classes.

There are a total of five dissimilarity measuring techniques, three clustering algorithms and two methods that wrap the distance and clustering together. The distance measures are the traditional daisy method, the newer random forest technique and our proposed distances d_1 , d_2 , and d_3 . The three clustering algorithms are Agnes, PAM and K -means. K -means is used only with numeric datasets Iris and optical digits. The two sparse methods, K -means sparse and hierarchical sparse, combine the distance calculation and clustering using techniques previously described.

B. RESULTS

This analysis was done on a MacBook Pro with a 2.6Ghz Intel Core i7 running 16GB of RAM. R Studio with R version 3.1.1 (R Core Team, 2014) was used execute the code. For each dataset, two sets of noise variables are added. Daisy dissimilarity is then computed for all datasets (for the numeric datasets Iris and optical digits, the daisy dissimilarity is Euclidean distance). Random forest proximities and d_1 through d_3 distances are also computed for all datasets. All were used in Agnes and PAM clustering algorithms. All of the techniques were computationally tractable, and the time to run each one was less than five minutes. Sparse K -means was run only on the two numerical datasets, Iris and optical digits, and took less than 10 minutes on each. Sparse hierarchical clustering was quick, except on optical digits, which took about an hour to run. The resulting Cramér's V for each run are displayed in the Tables 3–6.

For the Iris dataset shown in Table 3, d_2 produces the highest Cramér's V values (highlighted in yellow) independent of the clustering algorithm. The best clustering results are produced when d_2 is paired with PAM (highlighted in yellow). In the dataset with 15 noise variables and run using 2K clusters, Agnes produces a slightly better solution than PAM. Using 50 noise variables and 2K clusters, d_1 clustered with Agnes also produces a slightly better solution.

Table 3. Iris Dataset Results

Dataset	Clusters	Daisy			Sparse		Random Forest	
		Agnes	Pam	K Means	KMSparse	HeirSparse	Agnes	Pam
Iris	3	0.7075	0.8633	0.8633	0.8839	0.8452	0.8513	0.8360
Iris	6	0.7808	0.9261	0.9199	0.8984	0.8700	0.9529	0.8923
Iris 15 noise	3	0.1157	0.6526	0.7307	0.9081	0.8392	0.6451	0.4841
Iris 15 noise	6	0.6833	0.6643	0.6844	0.9147	0.9032	0.6625	0.5279
Iris 50 noise	3	0.1162	0.4205	0.6972	0.8839	0.7670	0.1769	0.1992
Iris 50 noise	6	0.2183	0.5024	0.7252	0.8787	0.9343	0.2923	0.3261

Dataset	Clusters	d_1		d_2		d_3	
		Agnes	Pam	Agnes	Pam	Agnes	Pam
Iris	3	0.8771	0.8909	0.8771	0.9081	0.8058	0.8058
Iris	6	0.9370	0.9316	0.9363	0.9458	0.9359	0.9440
Iris 15 noise	3	0.8058	0.8276	0.8276	0.9514	0.7223	0.8058
Iris 15 noise	6	0.9485	0.9441	0.9531	0.9444	0.8095	0.8631
Iris 50 noise	3	0.7217	0.7563	0.9333	0.9514	0.7071	0.7085
Iris 50 noise	6	0.9446	0.8922	0.9442	0.9322	0.7155	0.7608

The second numerical dataset in Table 4, optical digits, favors sparse K -means. The new distance metrics perform well, but sparse K -means beats them by a small margin. Even in the presence of noise, the sparse K -means algorithm maintained its advantage because it is able to minimize the contribution of variables that add no value to a dissimilarity calculation.

Table 4. Optical Digits Dataset Results

Dataset	Clusters	Daisy			Sparse		Random Forest	
		Agnes	Pam	K Means	KMSparse	HeirSparse	Agnes	Pam
Optical	10	0.7106	0.8116	0.7956	0.8217	0.3944	0.7892	0.7166
Optical	20	0.8534	0.8768	0.9123	0.9108	0.4749	0.8861	0.6759
Optical 15 noise	10	0.7005	0.7485	0.7822	0.8218	0.3887	0.6793	0.4804
Optical 15 noise	20	0.8472	0.8601	0.9081	0.9132	0.4897	0.8340	0.7928
Optical 50 noise	10	0.7757	0.6585	0.8203	0.8242	0.3638	0.5114	0.7821
Optical 50 noise	20	0.8047	0.7831	0.8872	0.9157	0.4755	0.5323	0.5548

Dataset	Clusters	d_1		d_2		d_3	
		Agnes	Pam	Agnes	Pam	Agnes	Pam
Optical	10	0.7526	0.7226	0.7128	0.7340	0.6936	0.7364
Optical	20	0.8354	0.7225	0.8391	0.7350	0.8440	0.7364
Optical 15 noise	10	0.7582	0.7311	0.6743	0.7352	0.7671	0.7346
Optical 15 noise	20	0.8614	0.8168	0.8396	0.8065	0.8470	0.8368
Optical 50 noise	10	0.7566	0.8136	0.7530	0.7940	0.6215	0.8368
Optical 50 noise	20	0.8502	0.8007	0.8435	0.7934	0.8285	0.8490

The Splice dataset in Table 5 is categorical, so the K -means and sparse K -means algorithms are not used. With this dataset, d_2 clearly outperformed other techniques. The introduction of noise brought small improvements, due to the automatic variable selection offered by the algorithm.

Table 5. Splice Dataset Results

Dataset	Clusters	Daisy			Sparse		Random Forest	
		Agnes	Pam	K Means	KMSparse	HeirSparse	Agnes	Pam
Splice	3	0.3600	0.1782			0.0617	0.0631	0.0893
Splice	6	0.3603	0.3038			0.0877	0.1206	0.1283
Splice 15 noise	3	0.2609	0.1786			0.0617	0.1994	0.1480
Splice 15 noise	6	0.2877	0.2638			0.0982	0.2578	0.1358
Splice 50 noise	3	0.2329	0.1600			0.0145	0.0668	0.0858
Splice 50 noise	6	0.2901	0.2353			0.0398	0.1515	0.1138

Dataset	Clusters	d_1		d_2		d_3	
		Agnes	Pam	Agnes	Pam	Agnes	Pam
Splice	3	0.5943	0.5017	0.5977	0.7815	0.0419	0.5647
Splice	6	0.5971	0.6040	0.8249	0.7988	0.0664	0.5952
Splice 15 noise	3	0.6104	0.5174	0.6347	0.7677	0.0359	0.6092
Splice 15 noise	6	0.6128	0.6130	0.8312	0.7982	0.5738	0.6358
Splice 50 noise	3	0.6104	0.5174	0.6347	0.7677	0.0359	0.6092
Splice 50 noise	6	0.6128	0.6130	0.8312	0.7982	0.5738	0.6358

Cardiac arrhythmia was the final dataset shown in Table 6. It contained mixed variable types and performed best with the tree distances. This was the

first dataset in which d_3 performed strongly. The metrics d_1 and d_2 did well in a noisier environment.

Table 6. Cardiac Arrhythmia Dataset Results

Dataset	Clusters	Daisy			Sparse		Random Forest	
		Agnes	Pam	K Means	KMSparse	HeirSparse	Agnes	Pam
Arrhythmia	13	0.3406	0.3877			0.3143	0.3750	0.3235
Arrhythmia	26	0.4998	0.5014			0.4032	0.4489	0.4344
Arrhythmia 15 noise	13	0.2898	0.3381			0.3238	0.4063	0.2777
Arrhythmia 15 noise	26	0.4160	0.4821			0.4059	0.4676	0.3975
Arrhythmia 50 noise	13	0.3276	0.3729			0.3355	0.2933	0.3048
Arrhythmia 50 noise	26	0.4880	0.4509			0.4220	0.3903	0.4015

Dataset	Clusters	d_1		d_2		d_3	
		Agnes	Pam	Agnes	Pam	Agnes	Pam
Arrhythmia	13	0.4436	0.3659	0.4231	0.3549	0.4502	0.3795
Arrhythmia	26	0.5060	0.4802	0.4939	0.4841	0.5179	0.4898
Arrhythmia 15 noise	13	0.4818	0.3934	0.4266	0.3681	0.4084	0.3063
Arrhythmia 15 noise	26	0.5255	0.4880	0.5461	0.4775	0.4928	0.4810
Arrhythmia 50 noise	13	0.4666	0.3837	0.4607	0.3225	0.4311	0.4072
Arrhythmia 50 noise	26	0.5156	0.4776	0.4990	0.4692	0.5172	0.4796

C. CONCLUSION

The Buttrey tree distances performed best with categorical and mixed datasets. They performed well with numerical data, but sparse K -means slightly outperformed them when dealing with the optical digits dataset. It was expected that as the new distance metrics increased in complexity, the resulting clustering solutions would improve. However, in this analysis the best result was the intermediate distance algorithm, d_2 .

V. SUMMARY

A. SUMMARY

This thesis examines new distance measurements that offer independence from variable scaling, resistance to monotonic transformations, automatic variable selection and are able to process datasets of mixed variable types. Chapter I introduced the problem and laid out the objectives. Chapter II's literature review was a background on past and current state of the clustering arts. Chapter III reviewed the tree distances and discussed the datasets to be used in this analysis. The fourth chapter presented the results of this study.

Of the three tree distances, d_2 consistently performed well and simultaneously dealt with the four problems addressed at the outset. Tree distances best handled categorical and mixed variable types, with d_2 performing best overall. Automatic variable selection was tested with the addition of noise variables, and tree distances performed well. Their nearest competitor was sparse clustering, which was computationally expensive with large datasets, and had difficulty with categorical data. The last two problems of variable scaling and monotonic transformations were dealt with by the use of CART upon which tree distances are based.

B. FUTURE WORK

It was expected that as the tree distances increased in complexity, their performance would improve. There was no significant improvement from d_2 to d_3 , so work remains to be done on whether evaluating deviance decreases at parent nodes can produce better results.

We discussed the increasing volume of data that organizations now deal with and will encounter in the future. Application of tree distance to “big data” and comparisons of computations tractability to these future clustering problems should be explored. Parallel processing or high power computing can be used with “big data” to explore processing speed, and also visualization tools.

Visualization tools are another important application of tree distances. Multi-dimensional scaling tools require a dissimilarity matrix to produce 2-D and 3-D images of high dimensional data. This can be used in addition to or instead of clustering to give users a better understanding underlying data structures and can benefit from the computational ease and relative accuracy of tree distance calculations.

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